PATENT Attorney Docket No. H 3491 PCT/US

IN THE UNITED STATES PATENT AND TRADEMARK OFFICE

CERTIFICATE OF EXPRESS MAIL (37 C.F R § 1 10)

I HEREBY CERTIFY THAT THIS PAPER OR FEE IS BEING DEPOSITED WITH THE UNITED STATES POSTAL SERVICE, EXPRESS MAIL POST OFFICE TO ADDRESSEE" UNDER 37 C.F.R. § 1.10, BEARING EXPRESS MAIL LABEL NO. EK219526757US ON THIS 1ST DAY OF DECEMBER, 2000 AND IS ADDRESSED TO: COMMISSIONER FOR PATENTS, WASHINGTON, D.C. 2023

Int'l Appl. No.: PCT/EP99/03563

Int'l Filing Date: May 25, 1999

Priority Date: June 3, 1998

Applicants: Beatrix KOTTWITZ et al.

Serial No.: To be assigned

Filed: December 1, 2000

For: DETERGENT CONTAINING AMYLASE

AND PERCARBONATE

AND PERCARBONATE

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Commissioner for Patents Box PCT Washington, D.C. 20231

TRANSMITTAL OF APPLICATION PAPERS TO U.S. DESIGNATED\ELECTED OFFICE (DO/EO/US) CONCERNING A FILING UNDER 35 U.S.C. § 371 (37 CFR 1.494 OR 1.495)

This transmittal letter is based upon Form PTO-1390.

The above-identified applicant has filed an International Application under the PCT and hereby submit(s) to the United States Designated/Elected Office (DO/EO/US) the following items and other information:

1. [X] This is the FIRST submission of items concerning a filing under 35 U.S.C. §371.

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International Application No. PCT/EP99/03563

| 2. | This is the SECOND or SUBSEQUENT submission of items concerning a filing |
|----|--|
| | under 35 U.S.C. §371. |

- This is an express request to begin national examination procedures (35 U.S.C. §371(f)) at any time rather than delay.
- 4. [X] A proper Demand for International Preliminary Examination (IPE) was made to the appropriate Authority (IPEA) by the 19th month from the earliest claimed priority date (see attached).
- 5. [X] A copy of the International Application as amended (35 U.S.C. §371(c)(2)) -
 - (a) [] is transmitted herewith (required when not transmitted by International Bureau).
 - (b) [X] has been transmitted by the International Bureau.
 - (c) [] is not required, as the application was filed in the United States Receiving Office (RO/US).
- 6. [X] A translation of the International Application into the English language is enclosed.
- 7. [X] Amendments to the (specification and) claims of the International Application under PCT Article 19 (35 U.S.C. §371 (c)(3))
 - (a) [] are transmitted herewith (required if not transmitted by the International Bureau).
 - (b) | have been transmitted by the International Bureau.
 - (c) [] have not been made; however, the time limit for making such amendments has NOT expired.
 - (d) [X] have not been made and will not be made.
 - (e) [] will be submitted with the appropriate surcharge.
- 8. [] A translation of the amendments to the claims (and/or the specification) under PCT Article 19 (35 U.S.C. §173(c)(3)) is enclosed or will be submitted with the appropriate surcharge.
- 9. [X] An unexecuted Oath or Declaration/Power of Attorney of the inventor(s) (35 U.S.C. §371(c)(4)) is enclosed.
- 10. [] A translation of at least the Annexes to the IPE Report under PCT Article 36 (35 U.S.C. §371(c)(5)) is enclosed.

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Items 11. to 16. below concern other document(s) or information included:

- 11. An Information Disclosure Statement under 37 CFR 1.97 and 1.98 is enclosed.
- 12. An Assignment is enclosed for recording. A separate cover sheet in compliance with 37 CFR 3.28 and 3.31. is included.
- A FIRST preliminary amendment is enclosed. IT IS REQUESTED THAT THE FILING FEES FOR THE CLAIMS BE CALCULATED AFTER THE CLAIM AMENDMENTS IN THE PRELIMINARY AMENDMENT HAVE BEEN ENTERED.
- 14. [] A substitute specification (including claims, abstract, drawing) is enclosed.
- 15. A change of Power of Attorney and/or address letter is enclosed.
- 16. [X] Other items of information:
 - [] This application is being filed pursuant to 37 CFR 1.494(c) or 1.495(c), and any missing parts will be filed before expiration of -
 - [] 22 months from the priority date under 37 CFR 1.494(c), or
 - 32 months from the priority date under 37 CFR 1.495(c).
 - [] The undersigned attorney is authorized by the International application and by the inventors to enter the **National Phase** pursuant to 37 CFR 1.494(c) or 1.495(c).

The following additional information relates to the International Application:

- [x] Receiving Office: EPO (originally filed in the German Patent Office)
- [] IPEA (if filing under 37 CFR 1.495): EPO
- [x] Priority claim(s) (35 USC §§ 119, 365): App. No. 198 24 688.9 Filed: June 3, 1998
- [x] A copy of the International Search Report is
 - [x] enclosed.
 - [x] attached to the copy of the International Application.
- [x] A copy of the Receiving Office Request Form is enclosed.

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International Application No. PCT/EP99/03563

The fee calculation is set forth below.

[X]

FEE CALCULATION

Kindly send us the official filing receipt.

The Commissioner is hereby authorized to charge <u>any</u> additional fees which may be required or to credit any overpayment to Deposit Account No. 03-2775. This is a "general authorization" under 37 CFR 1.25(b), except that no <u>automatic</u> debit of the issue upon allowance is authorized.

Respectfully submitted,

CONNOLLY BOVE LODGE & HUTZ LLP

Date: December 1, 2000

James M. Olsen

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PATENT Attorney Docket No. H 3491 PCT/US

IN THE UNITED STATES PATENT AND TRADEMARK OFFICE

| Commissioner for Patents Box PCT Westington D.C. 20221 | |
|--|------------------------|
| AMYLASE AND PERCARBONATE |) |
| For: DETERGENTS CONTAINING |) |
| Filed: December 1, 2000 |) |
| Serial No.: To be assigned |) Examiner: Unassigned |
| Applicants: Beatrix KOTTWITZ et al. |) Art Unit: Unassigned |
| Priority Date: June 3, 1998 |) |
| Int'l Filing Date: May 25, 1999 |) |
| Int'l Appl. No.: PCT/EP99/03563 |) |

Washington, D.C. 20231

Sir:

PRELIMINARY AMENDMENT

Prior to the examination of the above application, please amend this application as follows:

IN THE CLAIMS:

Please cancel claims 2-11, without prejudice or disclaimer of the subject matter thereof.

IN THE ABSTRACT:

Please add the attached Abstract of the Disclosure.

REMARKS

If there are any other fees due in connection with the filing of this response, please charge

the fees to our Deposit Account No. 03-2775. If a fee is required for an extension of time under 37 C.F.R. § 1.136 not accounted for above, such an extension is requested and the fee should also be charged to our Deposit Account.

Respectfully submitted,

CONNOLLY BOVE LODGE & HUTZ LLP

Dated: December 1, 2000

James M. Olsen

Reg. No. 40,408

ABSTRACT OF THE DISCLOSURE

An amylase-containing detergent that includes α -amylase from Bacillus amyloliquefaciens and alkali metal percarbonate in addition to customary ingredients compatible with such constituents.

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WO 99/63036

Detergents comprising amylase and percarbonate

The present invention relates to enzyme containing detergents comprising besides customary constituents an amylase from Bacillus amyloliquefaciens and a certain peroxidic bleach.

Laundry detergents, in addition to the ingredients vital to the laundering process, such as surfactants generally comprise and builder materials, constituents, which may be grouped together under the heading of wash auxiliaries and which embrace such different active substance groups as foam regulators, graying inhibitors, bleaches, bleach activators, auxiliaries Such transfer inhibitors. color the surfactant assist which include substances performance by means of enzymatic degradation of stains present on the textile. The same applies, mutatis mutandis, to cleaning products for hard surfaces. As which assist removing in proteases, the well as and the fat-cleaving lipases, particular proteins, importance in this context attaches to the amylases. Amylases have the function of facilitating the removal of starchy stains by means of catalytic hydrolysis of the starch polysaccharide, and have been used for this time in dishwashing long for a fairly purpose detergents, but also in detergents for use in textile laundering. In by far the great majority of cases the amylase involved has comprised a heat-stable amylase from Bacillus licheniformis, which is customary in commerce under the designation Termamyl®, for example. More recently, there has been increased use in such compositions of genetically manipulated amylases; that amylases whose amino acid sequence been has altered, using gene technology methods, in comparison to naturally occurring amylases. As well as increasing their capacity to perform, the objective of genetically enhance is essentially to modifying amylases stability of the enzyme, especially their stability to attack by oxidizing agents. One approach toward achieving this objective, which was proposed International Patent Application WO 94/18314, consists in removing particularly oxidation-susceptible amino acids, such as methionine, tryptophan, cysteine or tyrosine, from the amino acid sequence of the amylase, or replacing them by other amino acids more stable to similar approach oxidation. A is proposed International Patent Application WO 95/21247, which recommends replacing at least one methionine in the amylase amino acid sequence by an amino acid which is neither methionine nor cysteine.

Although such genetic modifications may lead to improved amylase stability under certain application conditions, they do not help to increase the contribution of the amylase to the wash or cleaning performance of corresponding compositions comprising the amylase.

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It has surprisingly now been found that the combination of a naturally occurring α -amylase with a certain peroxidic oxidizing agent leads to unexpectedly synergistic performance improvements when used in detergents.

The invention accordingly provides an amylase containing detergent which comprises α -amylase from Bacillus amyloliquefaciens and an alkali percarbonate, as well as customary ingredients compatible with such constituents.

The invention further provides for the use of such a combination for enhancing the cleaning performance of 35 detergents, especially with respect to starchy and/or colored stains when used in detergent solutions. especially aqueous detergent solutions. In context, cleaning performance with respect to colored stains is to be understood in its widest context and

embraces not only the bleaching of soil present on the textile and the bleaching of soil which is in the wash liquor, having become detached from the textile, but also the oxidative destruction of textile colors present in the wash liquor, having become detached from textiles under the wash conditions, before they are able to attach to different-colored textiles. Also, in the context of use in cleaning solutions for hard surfaces, this term refers both to the bleaching of soil present on the hard surface, especially tea, and to the bleaching of soil which is present in the dishwashing liquor, having become detached from the hard surface.

- α -Amylase from Bacillus amyloliquefaciens has been known for a long time, for example, from the U.S. Patent US 1 227 374. It is available commercially, for example, under the designation Amylase BAN®.
- A composition of the invention contains preferably from 20 0.001 mg to 0.5 mg, in particular from 0.02 mg to 0.3 mg, of amylolytic protein per gram of the overall composition. The protein concentration determined using known methods, such as the bicinchonic acid technique (BCA technique, Pierce Chemical Co., 25 Rockford, IL) or the Biuret technique (A.G. Gornall, C.S. Bardawill and M.M. David, J. Biol. Chem. 751-766, 1948).
- A composition of the invention comprises preferably up 30 to 50% by weight, in particular from 5% by weight to 30% by weight, of alkali metal percarbonate, percarbonate being particularly preferred. It can be prepared by known processes and, especially for use in particulate compositions, may if desired be compounded 35 in granular form and/or stabilized and/or coated, as known, for example, from International Applications WO 91/15423, WO 92/17400, WO 92/17404. WO 93/04159, WO 93/04982, WO 93/20007, WO 94/03553,

WO 94/05594, WO 94/14701, WO 94/14702, WO 94/24044, WO 95/02555, WO 95/02672, WO 95/06615, WO 95/15291 or WO 95/15292 or in European Patent Applications EP 0 459 625, EP 0 487 256, EP 0 567 140, EP 0 623 553, EP 0 592 969 or EP 0 748 764. For reasons of stability it is preferably used in the form of granules which have been prepared from and/or coated with the aid of alkaline earth metal sulfate, alkali metal sulfate, alkali metal silicate, alkaline earth metal halide, alkali metal halide, alkali metal carbonate, metal hydrogencarbonate, alkali metal phosphate, alkali metal borate, alkali metal perborate, boric acid, partially hydrated alumosilicate, carboxylic acids, dicarboxylic acids, polymers of unsaturated carboxylic and/or dicarboxylic acids, or mixtures of these. In one preferred embodiment it has a morphology index (MI), as defined in EP 0 451 893, of below 0.06.

In one preferred embodiment of compositions of the invention, the ratio of α -amylase to alkali metal percarbonate is in the range from 0.0001 mg to 0.1 mg, in particular from 0.001 mg to 0.01 mg, of amylolytic protein per % by weight of alkali metal percarbonate in the detergent.

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In addition to the active substance combination used in accordance with the invention, the detergents of the invention, which may be present as solids, especially pulverulent solids, in postcompacted particle form, as homogeneous solutions or suspensions may in principle comprise all known ingredients which are customary in such compositions. The compositions of the invention in particular may comprise builder substances, surfactants, additional bleaches based on organic and/or inorganic peroxygen compounds, bleach activators, water miscible organic solvents, additional enzymes, sequesterants, electrolytes, pH regulators, and further auxiliaries, such as optical brighteners, graying inhibitors, color transfer inhibitors, foam regulators, silver corrosion inhibitors, and also dyes and fragrances.

compositions of the invention may comprise a surfactant or or two more surfactants, suitable surfactants comprising in particular anionic surfactants, nonionic surfactants and mixtures thereof, but also cationic, zwitterionic and amphoteric surfactants.

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Suitable nonionic surfactants are, in particular, alkyl glycosides and ethoxylation and/or propoxylation products of alkyl glycosides or linear or branched alcohols having in each case 12 to 18 carbon atoms in the alkyl moiety and from 3 to 12, preferably from 4 to 10, alkyl ether groups. It is also possible to use corresponding ethoxylation and/or propoxylation products of N-alkylamines, vicinal diols, fatty acid esters and fatty acid amides, which in terms of the alkyl moiety correspond to the aforementioned chain alcohol derivatives, and also alkylphenols having 5 to 12 carbon atoms in the alkyl radical.

Nonionic surfactants used are preferably alkoxylated, 25 advantageously ethoxylated, especially alcohols having preferably 8 to 18 carbon atoms and on average from 1 to 12 mol of ethylene oxide (EO) per mole of alcohol, in which the alcohol radical may be linear or, preferably, methyl-branched in position 2 and/or may comprise linear and methyl-branched radicals 30 in a mixture, as are commonly present in oxo alcohol radicals. In particular, however, preference is given to alcohol ethoxylates containing linear radicals from alcohols of natural origin having 12 to 18 carbon atoms, e.g., from coconut, palm, tallow fatty or oleyl 35 alcohol and on average from 2 to 8 EO per mole of alcohol. Preferred ethoxylated alcohols include, example, C_{12-14} alcohols containing 3 EO or 4 EO, C_{9-11} alcohols containing 7 EO, C_{13-15} alcohols containing

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3 EO, 5 EO, 7 EO or 8 EO, C_{12-18} alcohols containing 3 EO, 5 EO or 7 EO, and mixtures thereof, such as mixtures of C_{12-14} alcohol containing 3 EO and C_{12-18} alcohol containing 7 EO. The stated degrees ethoxylation represent statistical mean values, which for a specific product may be an integer or a fraction. Preferred alcohol ethoxylates have a narrowed homolog distribution (narrow range ethoxylates, NREs). addition to these nonionic surfactants it is also possible to use fatty alcohols containing more than 12 EO. Examples thereof are (tallow) fatty alcohols containing 14 EO, 16 EO, 20 EO, 25 EO, 30 EO or 40 EO. Especially in detergents for use in machine dishwashing processes, it is common to use extremely low-foaming compounds. These include, preferably, $C_{12}-C_{18}$ polyethylene glycol-polypropylene glycol ethers having in each case up to 8 mol of ethylene oxide propylene oxide units in the molecule. However, it is also possible to use other nonionic surfactants which are known to be low-foaming, such as, for example, C_{12} - C_{18} alkyl polyethylene glycol-polybutylene glycol ethers having in each case up to 8 mol of ethylene oxide and butylene oxide units in the molecule, endgroup-capped alkyl polyalkylene glycol mixed ethers. Particular preference is also given to the hydroxylcontaining alkoxylated alcohols as are described in European Patent Application EP 0 300 305, referred to as hydroxy mixed ethers. The nonionic surfactants also include alkyl glycosides of the general formula $RO(G)_{x}$, where R is a primary straight-chain or methyl-branched aliphatic radical, especially an aliphatic radical methyl-branched in position 2, containing 8 to preferably 12 to 18, carbon atoms, and G represents a glycose unit having 5 or 6 carbon atoms, preferably glucose. The degree of oligomerization, Х, which indicates distribution of the monoglycosides oligoglycosides, is any desired number - which, as a variable to be determined analytically, may also be a fraction - between 1 and 10; preferably, x is from 1.2

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to 1.4. Further suitable surfactants are polyhydroxy fatty acid amides of the formula (II)

$$\begin{array}{c|c}
R^{2} \\
\hline
S & | \\
R^{1}-CO-N-[Z]
\end{array} (II),$$

where R¹CO is an aliphatic acyl radical having 6 to 22 carbon atoms, R² is hydrogen or an alkyl or hydroxyalkyl radical having 1 to 4 carbon atoms, and [Z] is a linear or branched polyhydroxyalkyl radical having 3 to 10 carbon atoms and from 3 to 10 hydroxyl groups. The polyhydroxy fatty acid amides are derived preferably from reducing sugars having 5 or 6 carbon atoms, especially glucose. The group of the polyhydroxy fatty acid amides also includes compounds of the formula (III)

$$R^4-O-R^5$$

$$\downarrow$$

$$R^3-CO-N-[Z]$$
(III),

where R^3 is a linear or branched alkyl or alkenyl radical having 7 to 12 carbon atoms, R^4 is a linear, branched or cyclic alkylene radical or an arylene radical having 2 to 8 carbon atoms and R^5 is a linear, branched or cyclic alkyl radical or an aryl radical or an oxyalkyl radical having 1 to 8 carbon preference being given to C_1 - C_4 alkyl radicals or phenyl radicals, and [Z] is a linear polyhydroxyalkyl radical whose alkyl chain is substituted by at least two hydroxyl groups, or alkoxylated, preferably ethoxylated or propoxylated, derivatives of said radical. Here too, [Z] is preferably obtained by reductive amination of a sugar such as glucose, fructose, maltose, galactose, mannose, or xylose. The N-alkoxy-N-aryloxy-substituted compounds may then be converted to the desired polyhydroxy fatty acid amides, example, accordance in with the teaching of Patent Application WO 95/07331 International by

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reaction with fatty acid methyl esters in the presence of an alkoxide as catalyst. A further class of nonionic surfactants used with preference, which are used either as sole nonionic surfactant or in combination with other nonionic surfactants, in particular together with alkoxylated fatty alcohols and/or alkyl glycosides, are alkoxylated, preferably ethoxylated, or ethoxylated and propoxylated, fatty acid alkyl esters, preferably carbon atoms in the alkyl 1 to 4 especially fatty acid methyl esters, as are described, example, in Japanese Patent Application JP 58/217598, or those prepared preferably by the process described in International Patent Application WO 90/13533. Nonionic surfactants of the amine oxide type, examples N-cocoalkyl-N, N-dimethylamine being oxide and N-tallowalkyl-N,N-dihydroxyethylamine oxide, and of the fatty acid alkanolamide type, may also be suitable. The amount of these nonionic surfactants is preferably not more than that of the ethoxylated fatty alcohols, in particular not more than half thereof. Further suitable surfactants include those known as gemini surfactants. This term is used generally to refer to those compounds which possess two hydrophilic groups per molecule. These groups are generally separated from one another as a result of what is known as a spacer. This spacer is generally a carbon chain, which should be long enough to give the hydrophilic groups a sufficient spacing to allow them to independently of one another. Surfactants of this kind are generally notable for an unusually low critical micelle concentration and the ability to reduce greatly the surface tension of water. In exceptional cases, however, the expression gemini surfactants is used to embrace not only dimeric but also, correspondingly, surfactants. trimeric Examples of suitable surfactants sulfated hydroxy mixed are ethers accordance with German Patent Application DE 43 21 022 or dimer alcohol bis- and trimer alcohol tris-sulfates and ether sulfates in accordance with German Patent

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Application DE 195 03 061. Endgroup-capped dimeric and trimeric mixed ethers in accordance with German Patent Application DE-A-195 13 391 are notable in particular their biand multifunctionality. Thus endgroup-capped surfactants possess good wetting properties and are low-foaming, so making particularly suitable for use in machine washing or cleaning processes. However, it is also possible to use gemini-polyhydroxy fatty acid amides or polypolyhydroxy fatty acid amides, as described in International Patent Applications WO 95/19953, WO 95/19954, and WO 95/19955.

Suitable anionic surfactants are, in particular, soaps those containing sulfate or sulfonate groups. Preferred surfactants of the sulfonate type are C_{9-13} alkylbenzenesulfonates, olefinsulfonates, mixtures alkenesulfonates of and hydroxyalkanesulfonates, and also disulfonates, as are obtained, for example, from C_{12-18} monoolefins having a terminal or internal double bond by sulfonating with gaseous sulfur trioxide followed by alkaline or acidic hydrolysis of sulfonation products. Also suitable alkanesulfonates, which are obtained from C_{12-18} alkanes, for example, by sulfochlorination or sulfoxidation with subsequent hydrolysis or neutralization, respectively. Likewise suitable, in addition, are the esters of α -sulfo fatty acids (ester sulfonates), e.g., α -sulfonated methyl esters of hydrogenated coconut, palm kernel or tallow fatty acids, which are prepared by $\alpha\text{-sulfonation}$ of the methyl esters of fatty acids of plant and/or animal origin having 8 to 20 carbon atoms in the fatty acid molecule, followed by neutralization to give water-soluble mono-salts. Preferably, comprise the α -sulfonated esters of hydrogenated coconut, palm, palm kernel or tallow fatty acids, it being possible as well for sulfonation products of unsaturated fatty acids, e.g. oleic acid, to be present in small amounts, preferably in amounts of not more than about 2 to 3% by weight. Particular preference is

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given to $lpha ext{-sulfo}$ fatty acid alkyl esters having an alkyl chain of not more than 4 carbon atoms in the ester group, examples being methyl esters, esters, propyl esters, and butyl esters. particular advantage, the methyl esters of the lpha-sulfo fatty acids (MES) are used, and also their saponified suitable anionic surfactants di-salts. Further sulfated fatty acid glycerol esters which are the diesters and triesters, monoesters, and mixtures thereof, as obtained in the preparation esterification of a monoglycerol with from 1 to 3 mol fatty acid or in the transesterification triglycerides with from 0.3 to 2 mol of glycerol. Preferred alk(en)yl sulfates are the alkali metal salts, and especially the sodium salts, of the sulfuric monoesters of $C_{12}\text{-}C_{18}$ fatty alcohols, examples being those of coconut fatty alcohol, tallow fatty alcohol, lauryl, myristyl, cetyl or stearyl alcohol, or of $C_{10}\mbox{-}C_{20}$ oxo alcohols, and those monoesters of secondary alcohols of this chain length. Preference is also given alk(en)yl sulfates of said chain length contain a synthetic straight-chain alkyl radical prepared on a petrochemical basis, these sulfates possessing degradation properties similar to those of the corresponding compounds based on fatty-chemical raw materials. From a detergents standpoint, $C_{12}\text{-}C_{16}$ alkyl sulfates and $\text{C}_{12}\text{-}\text{C}_{15}$ alkyl sulfates, and also $\text{C}_{14}\text{-}\text{C}_{15}$ alkyl sulfates, are particularly preferred. addition, 2,3-alkyl sulfates, which may for example be prepared in accordance with US Patents 3,234,258 or 5,075,041 and obtained as commercial products from Shell Oil Company under the name DAN®, are suitable anionic surfactants. Also suitable are the sulfuric monoesters of the straight-chain or branched C₇₋₂₁ alcohols ethoxylated with from 1 to 6 mol of ethylene oxide, such as 2-methyl-branched C_{9-11} alcohols containing on average 3.5 mol of ethylene oxide (EO) or fatty alcohols containing from 1 to Preferred anionic surfactants further include the salts

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of alkylsulfosuccinic acid, which are also referred to as sulfosuccinates or as sulfosuccinic esters and which constitute the monoesters and/or diesters sulfosuccinic acid with alcohols, preferably fatty alcohols and especially ethoxylated fatty alcohols. Preferred sulfosuccinates comprise C_{8-18} fatty alcohol radicals or mixtures thereof. Especially preferred sulfosuccinates contain a fatty alcohol radical derived ethoxylated fatty alcohols which themselves represent nonionic surfactants. Particular preference is given in turn to sulfosuccinates whose fatty alcohol radicals are derived from ethoxylated fatty alcohols having a narrowed homolog distribution. Similarly, it also possible use to alk(en)ylsuccinic containing preferably 8 to 18 carbon atoms alk(en)yl chain, or salts thereof. Further suitable anionic surfactants include fatty acid derivatives of amino acids, for example, of N-methyltaurine (taurides) of N-methylglycine (sarcosides). Particular preference is given here to sarcosides and to the sarcosinates and, of these, especially the sarcosinates higher fatty acids, which may be monopolyunsaturated, such as oleyl sarcosinate. Further suitable anionic surfactants are, in particular, soaps. Suitable soaps include in particular saturated fatty acid soaps, such as the salts of lauric acid, myristic acid, palmitic acid, stearic acid, hydrogenated erucic acid and behenic acid, and, in particular, mixtures of soaps derived from natural fatty acids, e.g., coconut, palm kernel, or tallow fatty acids. Together with these soaps, or as substitutes for soaps, it is also possible to use the known alkenylsuccinic salts.

The anionic surfactants, including the soaps, may be present in the form of their sodium, potassium or ammonium salts and also as soluble salts of organic bases, such as mono-, di- or triethanolamine. Preferably, the anionic surfactants are in the form of

their sodium or potassium salts, in particular in the form of the sodium salts.

In laundry detergents of the invention, surfactants are present in proportions of preferably from 5% by weight to 50% by weight, in particular from 8% by weight to 30% by weight, whereas compositions for cleaning hard surfaces, especially for the machine cleaning of kitchen- and tableware, have lower surfactant contents of up to 10% by weight, in particular up to 5% by weight, and preferably in the range from 0.5% by weight to 3% by weight.

A composition of the invention comprises preferably at least one water soluble and/or water insoluble, organic 15 and/or inorganic builder. The water soluble organic builder substances include polycarboxylic especially citric acid and sugar acids, monomeric and polymeric amino polycarboxylic acids, in particular methylglycinediacetic acid, nitrilotriacetic acid and 20 ethylenediaminetetraacetic acid, and also polyaspartic acid, polyphosphonic acids, especially aminotris-(methylenephosphonic acid), ethylenediaminetetrakis-(methylenephosphonic and acid) 1-hydroxyethane-1,1diphosphonic acid, polymeric hydroxy compounds such as 25 dextrin, and also polymeric (poly)carboxylic especially the polycarboxylates of European EP 0 625 992 or of International Patent Application WO 92/18542 or of European Patent EP 0 232 202, which 30 obtainable by oxidizing polysaccharides respectively, dextrins; polymeric acrylic methacrylic acids, maleic acids and copolymers thereof, which may also contain in copolymerized form small fractions of polymerizable substances without carboxylic acid functionality. The relative molecular 35 mass of the homopolymers of unsaturated carboxyclic acids is generally between 3000 and 200,000, that of the copolymers between 2000 and 200,000, preferably from 30,000 to 120,000, based in each case on free

acid. A particularly preferred acrylic acid-maleic acid copolymer has a relative molecular mass of from 30,000 to 100,000. Commercially customary products are, for example, Sokalan® CP 5, CP 10 and PA 30 from BASF. Suitable though less preferred compounds of this class 5 are copolymers of acrylic acid or methacrylic acid with vinyl ethers, such as vinyl methyl ethers, esters, ethylene, propylene and styrene, in which the acid fraction is at least 50% by weight. As water soluble organic builder substances it is also possible 10 terpolymers containing as monomers unsaturated acids and/or their salts and, as the third monomer, vinyl alcohol and/or an esterified alcohol or a carbohydrate. The first acidic monomer or 15 salt thereof is derived from a monoethylenically unsaturated C_3 - C_8 carboxylic acid and, preferably, from C_3-C_4 monocarboxylic acid, especially (meth)acrylic acid. The second acidic monomer or salt thereof may be a derivative of a C_4-C_8 dicarboxylic acid, maleic acid being particularly preferred, and/or 20 a derivative of an allylsulfonic acid substituted in position 2 by an alkyl or aryl radical. Such polymers may be prepared in particular by processes which are described in German Patent DE 42 21 381 and German Patent Application DE 43 00 772, and generally have a 25 relative molecular mass of between 1000 and 200,000. Further preferred copolymers are those described in German Patent Applications DE 43 03 320 DE 44 17 734, containing monomers as preferably acrolein and acrylic acid/acrylic acid salts, 30 acetate. The organic builder substances, especially for the preparation of liquid compositions, may be used in the form of aqueous solutions, preferably aqueous solutions with a strength of from 30 to 50 percent by weight. All said acids are generally 35 in form of their water the soluble especially their alkali metal salts.

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Organic builder substances of this kind may be present, if desired, in amounts of up to 40% by weight, in particular up to 25% by weight, and preferably from 1% by weight to 8% by weight. Amounts close to the stated upper limit are used preferably in pastelike or liquid compositions of the invention, especially those containing water.

soluble inorganic builder materials Suitable water include, in particular, alkali metal silicates, alkali metal carbonates and alkali metal phosphates, which may be present in the form of their alkaline, neutral or acidic sodium or potassium salts. Examples thereof are trisodium phosphate, tetrasodium diphosphate, disodium dihydrogen diphosphate, pentasodium triphosphate, called sodium hexametaphosphate, oligomeric trisodium phosphate having degrees of oligomerization from 5 to 1000, especially from 5 to 50, and also corresponding potassium salts and/or mixtures of sodium and potassium salts. Water insoluble, water dispersible inorganic builder materials used are, in particular, crystalline or amorphous alkali metal alumosilicates, in amounts of up to 50% by weight, preferably not more than 40% by weight, and in liquid compositions in particular from 1% by weight to 5% by weight. Among these, preference is given to the crystalline sodium alumosilicates of detergent quality, especially zeolite A, P and, if appropriate, X, alone or in mixtures, in the form for example of a cocrystallizate of zeolites A and X (Vegobond® AX, a commercial product from Condea Augusta S.p.A.). Amounts close to the stated upper used preferably limit in solid, particulate compositions. Suitable alumosilicates possess particular no particles having a size of more than 30 $\mu m\text{,}$ and preferably consist at least 80% by weight of particles having a size below 10 $\mu\text{m}\text{.}$ Their calcium binding capacity, which may be determined in accordance with the information in German Patent DE 24 12 837, is

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generally in the range from 100 to 200 mg of CaO per gram.

Suitable substitutes or partial substitutes for said alumosilicate are crystalline alkali metal silicates, which may be present alone or in a mixture amorphous silicates. The alkali metal silicates which can be used as builders in the compositions of the invention preferably have a molar ratio of alkali metal oxide to SiO_2 of below 0.95, in particular from 1:1.1 to 1:12, and may be present in amorphous or crystalline form. Preferred alkali metal silicates are the sodium siliates, especially the amorphous sodium silicates, having a molar ratio $Na_2O:SiO_2$ of from 1:2 to 1:2.8. Those with a molar ratio $Na_2O:SiO_2$ of from 1:1.9 to 1:2.8 may be prepared by the process of European Patent Application EP 0 425 427. As crystalline silicates which may be present alone or in a mixture with amorphous silicates it is preferred to use crystalline phyllosilicates of the general formula $Na_2Si_xO_{2x+1} \cdot yH_2O$, where x, the so-called modulus, is a number from 1.9 to 22, in particular from 1.9 to 4, and y is a number from 0 to 33, and preferred values for x are 2, 3 or 4. Crystalline phyllosilicates which fall under general formula are described, for example, in European Patent Application EP 0 164 514. Preferred crystalline phyllosilicates are those where x in the stated general formula adopts the values 2 or 3. In particular, both and δ -sodium disilicates $(Na_2Si_2O_5 \cdot yH_2O)$ preferred, with β -sodium disilicate, for example, being obtainable by the process described in International Application WO 91/08171. δ -Sodium silicates having a modulus of between 1.9 and 3.2 may be prepared accordance with Japanese Patent Applications JP 04/238 809 and JP 04/260 610. In addition, virtually anhydrous crystalline alkali metal silicates of the abovementioned general formula wherein x is a number from 1.9 to 2.1, prepared from amorphous alkali metal silicates, and preparable as described in European

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Patent Applications EP 0 548 599, EP 0 502 325 and EP 0 452 428, may be used in compositions the invention. In a further preferred embodiment compositions of the invention, a crystalline sodium phyllosilicate having a modulus of from 2 to 3 is used, as may be prepared from sand and soda by the process of European Patent Application EP 0 436 835. Crystalline sodium silicates having a modulus in the range from 1.9 to 3.5, as are obtainable by the processes of European Patents EP 0 164 552 and/or EP 0 294 753, are used in a further preferred embodiment of compositions of invention. Crystalline sheetlike silicates of the abovementioned formula are sold by Clariant GmbH under the trade name Na-SKS, e.g., Na-SKS-1 (Na $_2$ Si $_{22}$ O $_{45} \cdot xH_2$ O, kenyaite), Na-SKS-2 (Na₂Si₁₄O₂₉·xH₂O, magadiite), Na-SKS-3 (Na₂Si₈O₁₇·xH₂O) or Na-SKS-4 (Na₂Si₄O₉·xH₂O, makatite). Of these, those particularly suitable include Na-SKS-5 $(\alpha-Na_2Si_2O_5)$, Na-SKS-7 $(\beta-Na_2Si_2O_5)$, natrosilite), Na-SKS-9 (NaHSi₂O₅·3H₂O), Na-SKS-10 (NaHSi₂O₅·3H₂O, kanemite), Na-SKS-11 $(t-Na_2Si_2O_5)$ and Na-SKS-13 (NaHSi₂O₅), especially Na-SKS-6 $(\delta-Na_2Si_2O_5)$. Α review crystalline phyllosilicates is given, for example, by the articles published in "Hoechst High Chem Magazin 14/1993" on pages 33-38 and in "Seifen-Öle-Fette-Wachse, Vol. 116, No. 20/1990" on pages 805-808. In one preferred embodiment of compositions of the invention, is made of a granular compound of crystalline phyllosilicate and citrate, of crystalline phyllosilicate and abovementioned (co)polymeric polycarboxylic acid, as is described, for example, German Patent Application DE 198 19 187, or of alkali metal silicate and alkali metal carbonate, described, for example, in International Patent Application WO 95/22592 or as is available commercially, for example, under the name Nabion® 15.

Builder substances may be present in the compositions of the invention, if desired, in amounts of up to 90% by weight. They are preferably present in amounts of up

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to 75% by weight. Laundry detergents of the invention have builder contents of in particular from 5% weight to 50% by weight. In compositions of invention for cleaning hard surfaces, especially for the machine cleaning of kitchen- and tableware, amount of builder substances is in particular from 5% by weight to 88% by weight, preferably no insoluble builder materials being used in compositions. In a preferred embodiment of compositions of the invention intended in particular for the machine washing of kitchen- and tableware, from 20% by weight to 40% by weight of water soluble organic builder, especially alkali metal citrate, from 5% by weight to 15% by weight of alkali metal carbonate, and from 20% by weight to 40% by weight of alkali metal disilicate are present.

Additional peroxygen compounds that may be suitable for compositions in of the invention include, 20 particular, organic peracids or peracidic salts of organic acids, such as phthalimidopercaproic perbenzoic acid or salts of diperdodecanedioic acid, hydrogen peroxide, and inorganic salts which give off hydrogen peroxide under the washing conditions. 25 including perborate, percarbonate, persilicate and/or persulfate such as Caroate. Where solid peroxygen compounds are additionally to be used, they may be employed in the form of powders or granules, which may also have been coated in a manner which is known in 30 principle. The addition of small amounts of known stabilizers bleach such as, for example, of phosphonates, borates and/or metaborates and metasilicates, and also magnesium salts such as magnesium sulfate, may also be judicious here.

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Bleach activators which may be used are compounds which under perhydrolysis conditions give rise to aliphatic peroxo carboxylic acids having preferably 1 to 10 carbon atoms, in particular 2 to 4 carbon atoms, and/or

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substituted or unsubstituted perbenzoic acid. Suitable substances are those which carry O-acyl and/or N-acyl groups of the stated number of carbon atoms, and/or substituted or unsubstituted benzoyl groups. Preference is given to polyacylated alkylenediamines, especially tetraacetylethylenediamine acylated triazine (TAED), derivatives, especially 1,5-diacetyl-2,4-dioxohexahydro-1,3,5-triazine (DADHT), acylated qlycolurils, especially tetraacetylglycoluril (TAGU), N-acyl imides, especially N-nonanoylsuccinimide (NOSI), acylated phenolsulfonates, especially n-nonanovlisononanoyloxybenzenesulfonate (nor iso-NOBS), carboxylic anhydrides, especially phthalic anhydride, acylated polyhydric alcohols, especially triacetin, ethylene glycol diacetate, 2,5-diacetoxy-2,5dihydrofuran, and the enol esters known from German Patent Applications DE 196 16 693 and DE 196 16 767, and also acetylated sorbitol and mannitol and/or the mixtures thereof (SORMAN) described in European Patent Application EP 0 525 239, acylated sugar derivatives, especially pentaacetylglucose (PAG), pentaacetylfructose, tetraacetylxylose and octaacetyllactose, and acetylated, optionally N-alkylated glucamine gluconolactone, and/or N-acylated lactams, for example, N-benzoylcaprolactam, which known are from International Patent Applications WO 94/27970, WO 94/28102, WO 94/28103, WO 95/00626, WO 95/14759 and WO 95/17498. The hydrophilically substituted acylacetals known from German Patent Application DE 196 16 769 and acyllactams described in German Patent Application DE 196 16 770 and in International Patent Application WO 95/14075 are likewise used with The combinations of conventional bleach preference. activators known from German Patent Application DE 44 43 177, may also be used. Such bleach activators especially when the abovementioned peroxide donating bleaches are present, be present in customary quantities, preferably in amounts of from 0.5% by weight to 10% by weight, and in particular from

1% by weight to 8% by weight, based on overall composition, but are preferably all absent when the percarboxylic acid essential to the invention is used as the sole bleach.

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In addition to the conventional bleach activators or instead of them it is also possible for sulfonimines and/or bleach boosting transition metal salts transition or metal complexes, European Patents EP 0 446 982 and EP 0 453 003, to be present as so-called bleaching catalysts, preferably in an amount of up to 1% by weight, in particular from 0.001% by weight to 0.25% by weight.

15 Enzymes which may be used in the compositions in addition to the amylase that is essential to the include those invention from the class the proteases, lipases, cutinases, pullulanases, hemicellulases, cellulases, oxidases, laccases 20 peroxidases, and mixtures thereof. If desired, amylases other than the amylase essential to the invention may be present in addition to it. Particularly suitable active enzymatic substances are those obtained from fungi or bacteria, such as Bacillus subtilis, Bacillus licheniformis, Bacillus lentus, Streptomyces griseus, 25 Humicola lanuginosa, Humicola insolens, Pseudomonas pseudoalcaligenes, Pseudomonas cepacia orCoprinus cinereus. The amylase essential to the invention and the enzymes that may be used in addition may - as described, for example, in European Patent EP 0 564 476 30 or in International Patent Application WO 94/23005 - be adsorbed on carrier substances and/or embedded coating substances in order to protect them against premature inactivation. In the detergents of invention they are present preferably in amounts of up 35 to 5% by weight, in particular from 0.2% by weight to 4% by weight. Where the composition of the invention comprises protease, it preferably has a proteolytic activity in the range from about 100 PU/g to about

10,000 PU/g, in particular from 300 PU/g to 8000 PU/g. Where two or more enzymes are to be used in the composition of the invention, this may be accomplished by incorporating the two or more separate, or conventionally separately compounded enzymes, or by means of two or more enzymes compounded together in granules, as known, for example, from International Patent Applications WO 96/00772 or WO 96/00773.

10 The organic solvents which may be used besides water in the compositions of the invention, especially if they are present in liquid or paste form, include alcohols having 1 to 4 carbon atoms, especially methanol, ethanol, isopropanol and tert-butanol, diols having 2 15 to 4 carbon atoms, especially ethylene glycol propylene glycol, and also mixtures thereof and the ethers derivable from the aforementioned classes of compound. Water miscible solvents of this kind are present in the compositions of the invention preferably 20 amounts of not more than 30% by weight, particular from 6% by weight to 20% by weight.

Additionally, the compositions may comprise further constituents customary in detergents. These optional 25 constituents include, in particular, enzyme stabilizers, inhibitors, graying color transfer inhibitors, foam inhibitors, and optical brighteners and also dyes and fragrances. In order to protect against silver corrosion, silver corrosion inhibitors 30 may be used in dishwashing detergents of the invention. A cleaning product of the invention for hard surfaces may, furthermore, comprise abrasive constituents, especially from the group consisting of quartz flours, wood flours, polymer flours, chalks and microbeads, and mixtures thereof. Abrasives are present 35 in the detergents of the invention preferably at not more than 20% by weight, in particular from 5% by weight to 15% by weight.

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In order to adjust the pH to a desired level which does not come about of itself through the mixing of the other components, the compositions of the invention may and environmentally system-compatible comprise compatible acids, especially citric acid, acetic acid, tartaric acid, malic acid, lactic acid, glycolic acid, succinic acid, glutaric acid and/or adipic acid, or else mineral acids, especially sulfuric acid, or bases, ammonium hydroxides or alkali especially hydroxides. pH regulators of this kind are present in compositions of the invention in preferably not more than 20% by weight, in particular from 1.2% by weight to 17% by weight.

15 The color transfer inhibitors suitable for use in laundry detergents of the invention include, in particular, polyvinylpyrrolidones, polyvinylimidazoles, polymeric N-oxides such as poly(vinylpyridine N-oxide), and copolymers of vinylpyrrolidone with vinylimidazole.

Graying inhibitors have the function of keeping the soil detached from the textile fiber in suspension in the liquor. Suitable for this purpose are water soluble colloids, usually organic in nature, examples being starch, glue, gelatin, salts of ether carboxylic acids or ether sulfonic acids of starch or of cellulose, or salts of acidic sulfuric esters of cellulose or of Water soluble polyamides containing starch. groups are also suitable for this purpose. Furthermore, starch derivatives other than those mentioned above may be used, aldehyde starches, for example. Preference is cellulose to ethers, such as methylcellulose salt), methylcellulose, (Na hydroxyalkylcellulose and mixed ethers, such methylhydroxyethylcellulose, methylhydroxypropylcellulose, methylcarboxymethylcellulose, and mixtures thereof, in amounts, for example, of from 0.1 to 5% by weight, based on the compositions.

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As optical brighteners, laundry detergents of invention may comprise derivatives of diaminostilbenedisulfonic acid and/or its alkali metal Suitable, for example, are salts of 4,4'-bis(2-anilino-4-morpholino-1,3,5-triazinyl-6-amino)stilbene-2,2'disulfonic acid or compounds of similar structure which instead of the morpholino group carry a diethanolamino group, a methylamino group, an anilino group, or 2-methoxyethylamino group. It is also possible for brighteners of the substituted diphenylstyryl type to be present, for example, the alkali metal salts of 4,4'-bis(2-sulfostyryl)biphenyl, 4,4'-bis(4-chloro-3sulfostyryl) biphenyl, or 4-(4-chlorostyryl)-4'-(2sulfostyryl) biphenyl. Mixtures of the aforementioned optical brighteners may also be used.

Especially for use in machine processes, it may be of advantage to add customary foam inhibitors to the compositions. Examples of suitable foam inhibitors are soaps of natural or synthetic origin having a high $C_{18}-C_{24}$ fatty acid fraction. Examples of nonsurfactant-type foam inhibitors are organopolysiloxanes and their mixtures with microfine, optionally silanized silica and also paraffins, waxes, microcrystalline waxes, and mixtures thereof silanized silica or bis-fatty acid alkylene diamides. With advantages, use is also made of mixtures of different foam inhibitors, for example, mixtures comprising silicones, paraffins, or waxes. The foam inhibitors, especially those containing silicone and/or paraffin, are preferably bound on a granular, water soluble or dispersible support substance. preference given to mixtures of paraffins is bisstearylethylenediamide.

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The production of solid compositions of the invention presents no difficulties and may be done conventionally, for example, by spray drying or granulation, with the enzymes and any other heat

sensitive ingredients such as bleaches, for example, being added separately later on if desired. To produce compositions of the invention of heightened bulk density, in particular in the range from 650 g/l to 950 g/l, preference is given to a process known from European Patent EP 0 486 592 which includes an extrusion step. Another preferred preparation, with the aid of a granulation process, is described in European Patent EP 0 642 576.

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To produce compositions of the invention in tablet form, which may have one or more phases, may have one or more colors, and consist in particular of one layer or of two or more layers, in particular of two layers, it is preferred to follow a procedure in which all of the constituents - per one layer if appropriate - are mixed with one another in a mixer and the mixture is compressed by means of conventional tableting presses, for example, eccentric presses or rotary presses, at pressing forces in the range from about 50 to 100 kN, preferably from 60 to 70 kN. Especially in the case of multilayer tablets it may be of advantage if at least one layer is compressed beforehand. This is preferably accomplished at pressing forces of between 5 and 20 kN, in particular from 10 to 15 kN. In this way, tablets fracture resistant and yet dissolve sufficiently quickly under application conditions are obtained without problems, having fracture strengths and flexural strengths of normally from 100 to 200 N, but preferably above 150 N. A tablet produced in this way preferably has a weight of from 10 g to 50 g, in particular from 15 g to 40 g. The three-dimensional form of the tablets is arbitrary and may be circular, oval or angular, with intermediate forms also being possible. Corners and edges are advantageously rounded. Circular tablets preferably have a diameter of from 30 mm to 40 mm. In particular, the size of tablets of angular or cuboid design which are introduced predominantly by way of the dosing apparatus of, for

example, the dishwashing machine is dependent on the geometry and on the volume of this dosing apparatus. Embodiments which are preferred by way of example have a basal area of (from 20 to 30 mm) \times (from 34 to 40 mm), in particular of 26 \times 36 mm or of 24 \times 38 mm.

Liquid or paste detergents of the invention in the form of solutions comprising customary solvents are generally prepared by simply mixing the ingredients, which may be introduced in bulk or in solution into an automatic mixer.

Examples

15 Example 1

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To determine the washing power, cotton fabrics soiled with standardized test stains were washed at 40°C (detergent dosing 76 g; water hardness 16°d [German hardness]; load 3.5 kg, short program) in a domestic washing machine (Miele® W 701). Washing was carried out using a detergent **V1** containing 0.25% by weight of noninventive amylase granules (Termamyl® 60T) and 0.25% by weight of protease granules (activity 200,000 PE/g) containing the Bacillus lentus protease and 18% by weight of sodium percarbonate and 5% by weight of TAED, a detergent V2 of otherwise the same composition but containing, instead Termamyl®, of the proteinequivalent amount of the genetically modified amylase detergent **V3** of Duramyl®, a otherwise composition as V1 but containing, instead of Termamyl®, protein-equivalent amount of the genetically modified amylase Purafect® OxAm, a detergent V4 of otherwise the same composition as V1 but containing, instead of the Termamyl®, the protein-equivalent amount the fungal amylase Fungamyl®, and an inventive detergent M1 of otherwise the same composition as V1 but containing, instead of Termamyl®, the proteinequivalent amount of the amylase from Bacillus amyloliquefaciens.

The composition of the invention had a wash performance which was markedly superior to that of the compositions containing a different amylase. The situation is essentially the same when the percarbonate in the composition of the invention was replaced by perborate.

10 Example 2

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Detergents (V5 and V6, respectively) for the machine washing of kitchen- and tableware, comprising 55% by tripolyphosphate sodium (calculated anhydrous), 4% by weight amorphous sodium disilicate (calculated as anhydrous), 22% by weight carbonate, 9% by weight sodium percarbonate, 2% by weight TAED, 2% by weight nonionic surfactant, and also 1% or 2% by weight amylase granules (Termamyl® 60T) and by weight protease granules (activity 200,000 PU/g) comprising the Bacillus lentus protease (remainder to 100% by weight water, perfume and dye), (**V**7 and V8, compositions respectively), composition was otherwise like that of V5 and V6, respectively, but which instead of Termamyl® contained protein-equal amounts of Duramyl®, and compositions in accordance with the invention (M2 and M3, respectively), whose composition was otherwise like that of V5 and V6, respectively, but which instead of Termamyl® contained protein-equal amounts of amylase Bacillus amyloliquefaciens, were tested indicated below:

In a Miele® G 575 dishwasher (additions of 20 g of each test composition in the universal program, water hardness 14-16°dH [German hardness], operating temperature 55°C), in each case 6 plates soiled with standardized starch stains or bleachable stains were washed and the remaining stain residue was determined

gravimetrically and related to the initial value prior to washing (i.e., 100%). The compositions of significantly invention were superior to the noninventive compositions in cleaning performance. Essentially the same is true if the percarbonate in the compositions of the invention was replaced perborate.

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Annex to our letter of 05.29.2000 PCT/EP99/03563 (H 3491 PCT)

Replacement page

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What is claimed is:

- 1. An amylase containing detergent, charcterized in that it comprises α -amylase from Bacillus amyloliquefaciens and akali metal percarbonate in addition to customary ingredients compatible with such constituents, and does not comprise any antibody directed against the amylase.
- 10 2. The composition as claimed in claim 1, characterized in that it contains amylase amounts of from 0.001 mg to 0.5 mg, in particular from 0.02 mg to 0.3 mg, per gram of the overall composition.
 - 3. The composition as claimed in claim 1 or 2, characterized in that it contains up to 50% by weight, in particular from 5% by weight to 30% by weight, of alkali metal percarbonate.
 - 4. The composition as claimed in any of claims 1 to 3, characterized in that it contains from 0.5% by weight to 10% by weight, in particular from 1% by weight to 8% by weight, of compound which under perhydrolysis conditions gives off peoxocarboxylic acid.
- 5. The composition as claimed in any of claims 1 to 4, characterized in that it contains bleach boosting transition metal salts or transition metal complexes in an amount of up to 1% by weight, in particular from 0.001% by weight to 0.25% by weight.

- 6. The composition as claimed in any of claims 1 to 5, characterized in that the alkali metal percarbonate comprises sodium percarbonate.
- 5 7. The composition as claimed in any of claims 1 to 6, characterized in that the alkali percarbonate is used in the form of granules prepared and/or coated with the aid of alkaline earth metal sulfate, alkali metal sulfate, alkali 10 silicate, alkaline metal earth metal halide, alkali metal halide, alkali metal carbonate, alkali metal hydrogencarbonate, alkali phosphate, alkali metal borate, alkali metal perborate, boric acid, partially hydrated 15 alumosilicate, carboxylic acids, dicarboxylic acids, polymers of unsaturated carboxylic and/or dicarboxylic acids, or mixtures of these.
 - 8. The composition as claimed in any of claims 1 to 7, characterized in that the alkali metal percarbonate has a morphology index of below 0.06.
- 9. The composition as claimed in any of claims 1 to 8, characterized in that it contains from 0.0001 mg to 0.1 mg, in particular from 0.001 mg to 0.01 mg, of α -amylase per percent by weight of alkali metal percarbonate.
- 10. The use of a combination of α -amylase from 30 Bacillus amyloliquefaciens and alkali metal percarbonate for enhancing the cleaning performance of detergents which do not comprise any antibody directed against the amylase, when used in washing and cleaning solutions, especially 35 aqueous washing and cleaning solutions.

Replacement page

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11. The use as claimed in claim 10, characterized in that the cleaning performance is enhanced in relation to starchy and/or colored stains.

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| DETERGENT CONTAINING AMYLASE AND PERCARBONATE (Title of the Invention) is attached hereto OR | | | | | | | | | | | | |
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| City 40593 Duesseldorf | State | Z | Zip | Country | Germany | | Applicant | | | 1 |

| | price origin (| +) inside this box | | - | | | | | | | 1 3491 | PCT/US |
|--|------------------|-------------------------------|-------------|----------------|----------|-----|----------------------------|-----------------------------------|-------------------|----------------------------|--------------------|----------------|
| • | ` | DECL | .ARA1 | ΓΙΟΝ | | | | ADDITIC Sup | ONAL II piemen | NVENTOR Ital Sheet | (S) | |
| Name of Additional Joint Inventor, if any: A petition has been filed for this unsigned inventor | | | | | | | | | | | | ventor |
| Given Name | Ho | rst-Dieter | | Mid Initi | | | Family Name | SPECKMA | | | Suffix e.g. Jr. | Cittor |
| Inventor Signatur | 's H | out ai | bu to | red | a. | | | | Date | Dec. | L | 2000 |
| Residen City | ice: | Langenfeld | | St | ate | | Country | Germany | 4 | Citizenship | Germ | |
| ost Off | fice Addres | Eichenfeldst | rasse 24 a | | | | | | | <u> </u> | <u></u> | |
| ost Off | fice Addres | is | | | | | | | | | | |
| City | 40764 La | ngenfeld | State | | Zip | | Country | Germany | | Applicant Authority | | |
| lame | of Additi | onal Joint Inver | itor, if an | y: | | | A petition | has been fi | | | ned inv | /entor |
| Given Name | Kar | I-Heinz | | Mid Initi | | | Family Name | MAURER | • | | Suffix e.g. Jr. | |
| nventor': Signature | - 1 | Ral- de | 5 Mg | lu | 2 | 7 | | | Date | Dec. | | 2000 |
| Residenc City | ice: | Erkrath | . | Sta | ate | | Country | Germany | | Citizenship | Germ | |
| ost Offi | fice Addres | s Dechanstras | se 5 | | <u>.</u> | | | | <u></u> | | | |
| ost Offi | fice Addres | s | | | | | | | | | | |
| ity | 40699 Eri | krath | State | | Zip | | Country | Germany | | Applicant | | :- |
| lame | of Additi | onal Joint Inven | tor, if any | <i>/</i> : | | | petition | has been fi | | Authority this upsic | ned inv | ontor |
| Siven Iame | Chr | istian | | Mide Initia | | | Family Name | NITSCH | 100 101 | Ī | Suffix e.g. Jr. | entor |
| nventor's Signature | | 1/6:67 | \ | / | Z. | | 7 | | Date | Dec. | | 2000 |
| Residence | ce: | Duesseldorf | | Sta | ite | | Country | Germany | <u> </u> | Citizenship | Germa | |
| ost Offic | ice Addres | s Otto-Hahn-St | rasse 185 | | | | | | | | | <u></u> |
| | - 4 1 1 | s | | | | | | · · · · · · · · · · · · · · · · · | | | | |
| ost Offic | ice Addres | | | | | | | | | | | |
| | 40591 Du | esseldorf | State | | Zip | | Country | Germany | | Applicant | | |
| ity | 40591 Du | esseldorf onal Joint Inven | | · | Zip | Па | | | | Authority | and inv | onto: |
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| lame of the lame o | of Addition | | | Midd | lie | □ A | petition Family | | ed for | Authority this unsign | Suffix | entor |
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| lame of the lame o | of Additions e | onal Joint Inven | | Midd Initia | lie | | petition Family Name | | ed for | Authority this unsign | Suffix | entor |

Attorney Docket Number: H 3491 First Named Inventor: Beatrix

Beatrix KOTTWITZ

Title: DETERGENT CONTAINING AMYLASE AND

PERCARBONATE

Supplemental Sheet of Additional Attorney(s) and/or Agent(s):

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